In the Specification:

Please insert the following paragraph at the top of the first page and immediately before "CROSS-REFERENCE TO RELATED APPLICATIONS":

TITLE OF THE INVENTION

Carrier Protein Having an Adjuvant Effect

Please replace the paragraph immediately after "CROSS-REFERENCE TO RELATED APPLICATIONS" with the following:

This application is a division of Application Serial No. 09/679,750, filed 10/05/2000, now US Patent No. 6,780,420, which is a continuation of application Serial No. 08/836,500, filed 08/11/97, now US Patent No. 6,197,929, which is a national stage 371 application of the international application PCT/FR95/01463, which claims foreign priority to application 94,13306 filed 07/11/1994 in France.

Please insert the following heading immediately after the paragraph dealing with "CROSS-REFERENCE TO RELATED APPLICATIONS":

STATEMENT REGARDING FEDERALLY SPONSORED RESEARCH AND DEVELOPMENT

Not Applicable

Please insert the following heading immediately after the paragraph dealing with "STATEMENT REGARDING FEDERALLY SPONSORED RESEARCH AND DEVELOPMENT":

THE NAMES OF THE PARTIES TO A JOINT RESEARCH AGREEMENT

Not Applicable

Please insert the following heading immediately after the paragraph dealing with "THE NAMES OF THE PARTIES TO A JOINT RESEARCH AGREEMENT":

REFERENCE TO A "SEQUENCE LISTING"

The application contains "Sequence Listing" in the computer readable form. The computer readable form is identical with that filed in Application Serial No. 08/836,500, filed 08/11/97, now US Patent No. 6,197,929.

Please insert the following heading immediately after the paragraph dealing with "Reference to a "Sequence Listing":

FIELD OF THE INVENTION

Please insert the following heading immediately after line 9 of page 1 and immediately before line 10 (i.e. the sentence that starts, "The development of vaccines...") of page 1 in the originally submitted translation:

BACKGROUND ART

Please insert the following heading immediately after line 32 of page 1 and immediately before line 33 (i.e. the sentence that starts, "The Applicant has demonstrated...") of page 1 in the originally submitted translation:

SHORT SUMMARY OF THE INVENTION

Please insert the following heading immediately after line 11 of page 6 and immediately before line 12 (i.e. the sentence that starts, "In these examples...") of page 6 in the originally submitted translation:

BRIEF DESCRIPTION OF THE SEVERAL VIEWS OF THE DRAWINGS

Please insert the following heading immediately after line 24 of page 6 and immediately before line 25 (i.e. the sentence that starts, "Example 1: Isolation...") of page 6 in the originally submitted translation:

DETAILED DESCRIPTION OF THE INVENTION

Please replace the paragraph that starts on line 4 of page 7 with the following:

The pellets obtained after the second precipitation are resuspended in a 1% solution of zwittergent Zwittergent® 3-14.

Please replace the paragraph that starts on line 13, page 7 with the following:

The proteins of the MP fraction are dialysed against a 20 mM Tris/HC1, pH 8.0; 0.1% zwittergent Zwittergent © 3-14 buffer. The dialysate is loaded onto a column

containing a support of the strong anion exchanger type (column of $\emptyset = 50$ mm x H = 250 mm, Biorad® Macroprep High Q gel) which is equilibrated in the above-described buffer. The P40 protein is eluted at an NaCl concentration of 50 mM in the equilibration buffer.

Please replace the paragraph that starts on line 21, page 7 with the following:

The fractions containing the P40 are pooled and dialysed against a 20 mM citrate, pH 3.0; 0.1% <u>zwittergent @ 3-14</u> buffer. The dialysate is loaded onto a column containing a support of the strong cation exchanger type (diamensions of the column: $\emptyset = 25$ mm x H = 160 mm, Biorad@ Macroprep High S gel) which is equilibrated in the 20 mM citrate, pH 3.0; 0.1% <u>zwittergent Zwittergent @ 3-14</u> buffer. The P40 protein is eluted at an NaCl concentration of 0.7 M. The fractions containing the P40 are pooled and concentrated by ultrafiltration using a Minitan@ Millipore tangential flow filtration system employing membrane discs having a cutoff threshold of 10 kDa.

Please replace the paragraph that starts on line 9, page 10 with the following:

Gene amplification

Lysis buffer:

25 mM Taps, pH 9.3

2 mM MgCl₂

Amplification

buffer:

25 mM Taps, pH 9.3

2 mM MgCl₂ 0.1% Tween<u>®</u> 20 200 mM dNTP.

Please replace the paragraph that starts on line 16, page 10 with the following:

TST(20x):

Tris base 0.5 M

HC1 0.3 M

NaC1 4 M

Tween<u>®</u> 20 1%

EDTA 20 mM

Washing buffer:	Tris HC1	50 mM	pH 8.5
	$MgCl_2$	5 mM	
Denaturation	Gua – HC1	7.8 M	
solution:	Tris-HC1	28 mM	pH 8.5
Renaturation	Gua-HC1	0.5 M	
solution:	Tris-HC1	25 mM	pH 8.5
	NaC1	150 mM	
	Tween <u>®</u> 20	0.05%	

Please replace the paragraph that starts on line 24, page 11 with the following:

These reactions are carried out in 100 µ1 of amplification buffer using 5 pmol of each primer and 1 unit of Taq polymerase enzyme (Perkin Elmer Cetus). Each cycle comprises one denaturation step of 30 seconds at 95°C, followed by hybridization of the primer to the DNA and an extension of one minute at 72°C. 30 cycles are performed in this way using a Perkin Elmer Cetus "Gen Amp PCR" 9000 thermocycler.

Please replace the paragraph that starts on line 2, page 12 with the following:

The fragments which have thus been cloned are sequenced on an Applied Biosystems 373 automated DNA Sequencer. The sequencing reactions are carried out using the "Dye Terminator" kit in accordance with the supplier's (Applied Biosystems) recommendations either on double-stranded DNA obtained after gene amplification or derived from a maxiprep, or on single-stranded DNA drived from denatured PCR fragments (Hultman, T. et al., 1989, Nucleid Acids Rev. 17: 4937-4946).